ш



Europäisches Patentamt

European Patent Office

Office europé n des brevets



1) Publication number:

0 285 179 B1

(12)

EUROPEAN PATENT SPECIFICATION

- (5) Date of publication of patent specification: **16.06.93** (5) Int. CI.⁵: **G01N 33/533**, G01N 33/52, C07D 491/14
- (21) Application number: 88105319.3
- 2 Date of filing: 01.04.88

- Fluorescent diagnostic assay, assay elements, and conjugates.
- Priority: 02.04.87 US 34225
- Date of publication of application: 05.10.88 Bulletin 88/40
- Publication of the grant of the patent: 16.06.93 Bulletin 93/24
- Designated Contracting States:
 AT BE CH DE ES FR GB GR IT LI LU NL SE
- EP-A- 0 050 684 US-A- 4 622 400

JOURNAL OF IMMUNOLOGICAL METHODS, vol. 50, 1892, NL; J.A.TITUS et al., pp. 193-204

CLINICAL CHEMISTRY, vol. 31, no. 3, March 1985, Winston-Salem, NC (US); I.HEMMILÄ, pp. 359-370

ANALYTICAL CHEMISTRY, vol. 55, no. 4, April 1983, Washington, DC (US); B.WALTER, pp. 498a-514a

- 73 Proprietor: PB Diagnostic Systems, Inc. 151 University Avenue Westwood, MA 02090-2399(US)
- (2) Inventor: Arnost, Michael J. 29 Paddock Lane North Andover, MA 01845(US) inventor: Meneghini, Frank A. 123 Claremont Avenue Arlington, MA 02174(US) Inventor: Palumbo, Paul S. 271 Chesrry Street West Newton, MA 0216(US) Inventor: Stroud, Stephen G. 15 Dunbar Avenue Medford, MA 02155(US)
- Representative: Reitzner, Bruno, Dr. et al Patentanwälte Dipl.-Ing. R. Splanemann Dr. B. Reitzner, Dipl.-Ing. K. Baronetzky Tal 13 W-8000 München 2 (DE)

Note: Within nine months from the publication of the mention of the grant of the European patent, any person may give notice to the European Patent Office of opposition to the European patent granted. Notice of opposition shall be filed in a written reasoned statement. It shall not be deemed to have been filed until the opposition fee has been paid (Art. 99(1) European patent convention).

CHEMICAL ABSTRACTS, vol. 107, no. 7, 17 August 1987, Columbus, OH (US); M.C.GARNETT tal., p. 564, no. 57037z

CHEMICAL ABSTRACTS, vol. 100, no. 11, 12 March 1984, Columbus, OH (US); M.J.GEISOW et al., p. 262, no. 82277r

IMMUNOCHEMISTRY by L.A.Sternberger, Wiley Medical Publications, 3rd ed., 1986; pp. 44-47

Description

BACKGROUND OF THE INVENTION

It is known in the biological diagnostic assay field to utilize conjugates of biologically active moieties with a detectable, signal-generating dye moiety which may be, for example, a moiety which emits electromagnetic radiation, e.g., a fluorescent, chemiluminescent or bioluminescent moiety. The biologically active moiety may be: a DNA probe, e.g., a labeled DNA probe of the type used in detecting complementary DNA sequences; an enzyme; an enzyme inhibitor; an antigen; an antibody; a hapten, etc.

In recent years much attention has been focused on labeled-reagent immunoassays for the detection of body fluid components such as antigens, hormones, infectious agents, serum antibodies and the like. Consequently, the patent literature includes many disclosures of various assays involving a labeled-reagent reaction between antigens and antibodies to provide a detectable signal which may be a change in color, emission of electromagnetic radiation, etc. These assays involve an immunological interaction between a ligand and an antiligand wherein at least one of the two reactants contains a substance or a precursor of a substance which can provide the detectable signal as a function of the immunological ligand - antiligand interaction.

One class of labels commonly used in such assays are fluorescent dyes or fluorophores. Both heterogeneous and homogeneous specific binding assays employing fluorescent-labeled conjugates are well known in the art.

US-A-4 622 400 teaches certain rhodamine compounds which can be used as fluorescent tags in diagnostic methods. There is no suggestion that hydrophilic solubilizing groups can be incorporated in the dye compound.

EP-A-0 050 684 also teaches providing conjugates which includes a fluorescent dye attached to a biologically active moiety. There is no suggestion that a solubizing group is incorporated in the fluorescent dye molecule.

In general, it is desirable that fluorescent labels for use in such assays have a relatively long emission wavelength, e.g., above 500 mm. In addition, it is desirable that the labels have a large Stokes shift, be stable under the assay conditions, be relatively free of non-specific interference both from materials in solution and the moiety to which the label is conjugated and provide high quantum yields.

According to a first aspect, the present invention relates to a fluorescent diagnostic assay comprising combining in an assay medium: a sample suspected of containing an analyte; a binding partner for said analyte; and a conjugate of said analyte and a fluorescent compound, said conjugate represented by the formula

50

wherein W, X, Y and Z each independently is $\{CH_2\}_{a}(CHR_4)$ ($CH_2\}_{b}$;

R, R₁, R₃ and R₄ each independently is hydrogen, a substituent containing a hydrophilic group or a substituent containing a biologically active moiety; a

R₂ is -CO₂⁻, SO₃⁻ or a substituent containing a biologically active moiety;

provided that one of R_1 , R_2 , R_3 and R_4 is a substituent containing a biologically active moiety and at least one of the remaining of R_1 , R_3 and R_4 is a substituent containing a hydrophilic group;

EP 0 285 179 B1

a and b each independently is an integ r of from 0 to 4, provided that the sum of a + b is an integer of from 1 to 4; m and n each is 0 or 1 provided that on of m and n is 0 and the other is 1;

p and q each is 0 or 1 provided that one of p and q is 0 and th oth r is 1;

A is a counterion or the counterions to balance the overall charges on the conjugate moiety; and X is 0 or 1; separating bound conjugate from free conjugate; measuring the level of the fluorescent signal of said bound or free conjugate; and

relating said level of said signal to the amount of said analyte in said sample.

Under another aspect, the present invention relates to a diagnostic assay element adapted to receive a sample of a biological fluid and to provide a detectable signal as a function of an analyte which may be present in said fluid, said assay element containing a fluorescent conjugate represented by the general formula shown above.

Under a third aspect, the present invention relates to a fluorescent conjugate represented by general formula shown above.

Under a fourth aspect, the present invention relates to the use of the fluorescent conjugate as defined in the general formula above, for fluorescent staining of cells, or for the deletion or separation in a fluorescent activated cell sorter.

If R₁, R₃ and R₄ represent substituents containing a hydrophilic group, such group may be, for example, a carboxylic acid, a polyalcohol, a sulfonic acid or the like; if R, R₁, R₃ and R₄ represent a substituent containing a biologically active moiety, such moiety may be an antigen, an antibody or the like; this also applies for the biologically active moiety of R₂

It will be understood that where the conjugate moiety is neutral overall no counterion is needed. The counterion, A, may be any biologically acceptable anion or cation such as, for example, chloride, sulfate, diphenylphosphate, trifluoroacetate, trimethylammonium, sodium, potassium, calcium and the like. It will also be understood that when the conjugate moiety has a net overall charge greater than one, the charges may be balanced by one or more counterions. For example, where the conjugate moiety has a net overall charge of +2, these charges can be balanced by two chloride ions or one sulfate ion.

It should be noted that by "hydrophilic group", as used in the specification and claims herein, is meant a group which will improve the solubility of the molecule in water. Typical suitable hydrophilic groups which may be incorporated in the novel fluorescent compounds of the invention include: carboxylic acids (-COOH); polyethers such as those represented by

55 Et where c is an integer of from 1 to 20 such as polyethylene oxide; polyalcohols which are represented by

where d is an integer of from 1 to 20; primary, secondary or tertiary amines which are represented by -NR₅R₆ where R₅ and R₆ each independently is hydrogen, alkyl, preferably having from 1 to 6 carbon atoms, aryl such as phenyl or polyamines such as

$$-(CH_2)_2-N$$
 $-(CH_2)_2NH_2;$

sulfonic acids (-SO₃H); phosphonic acids or esters which are represented by

45

50

$$-(CH_2)_e$$
-PO(OR₇)(OR₈)

wher e is an integer of from 1 to 8 and R₂ and R₃ each independently is hydrogen, alkyl, preferably baving from 1 to 6 carbon atoms or aryl such as phenyl; phosphates represented by

$$-(CH2)e-OPO(OH)2;$$

phosphate esters represented by

10

15

30

35

$-(CH_2)_e$ OPO(OH)(OR₉)

where R₉ is alkyl, preferably having from 1 to 6 carbon atoms or aryl such as phenyl; phosphinic acids represented by

-(CH₂)_e-PO(OH)R₁₀

where R₁₀ is alkyl, preferably having from 1 to 6 carbon atoms; boronic acids which are represented by -R₁₁-B(OH)₂ where R₁₁ is

-(CH₂)e

or aryl such as phenyl; and borinic acids represented by -R₁₁-B(OH)R₁₀. As will be shown in more detail below, the compounds of the invention may contain one or more of the same hydrophilic groups or they may contain more than one different type of hydrophilic group.

The novel fluorescent conjugates within Formula I have emission maxima which are above 500 nm, typically in the range of from about 500 nm to about 650 nm.

Generally, the novel conjugates of the invention include at least one biologically active moiety which can be attached directly to the dye chromophore or which can be attached to the dye chromophore through a divalent achromophoric linking group. By the term "achromophoric linking group" is meant one which does not cause any appreciable shift in the spectral absorption characteristics of the dye moiety. Thus, in Formula I, where any of R, R₁, R₂, R₃ or R₄ is a substituent containing a biologically active moiety, such substituent may include a divalent achromophoric linking group to link the dye moiety to the biologically active moiety. Such a linkage should be non-conjugated.

The biologically active moiety may be any such as, for example, an antigen, an antibody, a hapten, a DNA probe, an Fab fragment.

Typical functional groups which are useful as coupling groups which can be attached to the dye molecule, and the substituents of the biologically active moiety with which such coupling groups are reactable to provide the achromophoric linking group within the labeled conjugates, are:

COUPLING GROUP

SUBSTITUENT OF BIOLOGICALLY ACTIVE GROUP

40	N-hydroxysuccinimide esters	amino groups (α -amino, lysine)
45	Imidoester	amino groups
	Aldehydes	amino groups
	Mixed anhydrides	nucleophilic groups
	Isothiocyanates	nucleophilic groups
	2,4-dichloro-5-triazine	nucleophilic groups
	Diazonium salts	tryptamine, histamine
50	Bromoacetyl	histamine, SH
	Maleimido	SH
	Activated disulfide bonds (e.g. 2-pyridyldisulfides)	SH

Many other linking groups may be incorporated into the conjugates of the invention. For xample, th hydrophilic groups mentioned above are substantially achromophoric and certain of these can be de-

EP 0 285 179 B1

rivatized and utilized as suitable linking groups. For example, an -NH₂ group can be converted to an amide as a result of attaching the biologically active moiety (BIO) to it, i.e., -NHCO-BIO. Further, in another embodiment a hydrophilic group may be attached to the linking group. For example, in the case of a primary amine, one of the hydrogen atoms is replaced by a biologically active moiety attached to it as described above and the other hydrogen atom can be replaced by a hydrophilic group such as, for example, -(CH₂)₂PO₃H₂. Thus, it will be appreciated that the linking group, while serving as the means for attaching the dye moiety to the biologically active moiety, can also have a hydrophilic group attached to it. In addition, the linking group may also function as a hydrophilic group such as, for example, in the case of -NH-BIO and -PO₂(OH)-BIO.

The labeled fluorescent conjugates of the invention are useful in various applications including diagnostic assays which are based on an energy transfer mechanism to activate the fluorescent label. The fluorescent dye moieties typically have maximum absorption, λ max, of from about 500 nm to about 650 nm, exhibit Stokes shifts of about 15 - 20 nm and have high quantum yields of about 0.7 - 0.8. These dye moieties advantageously offer a number of positions of attachment for biologically active moieties and/or solubilizing groups. The presence of such solubilizing groups can help to avoid undesirable nonspecific binding of the conjugates to components present in biological fluids such as, for example, plasma proteins.

DESCRIPTION OF THE PREFERRED EMBODIMENT

A prefered class of conjugates according to the invention which is used in preferred embodiments of the diagnostic assay and assay elements according to the invention, is represented by the formula

wherein R_{12} , R_{13} , R_{14} and R_{15} each independently is hydrogen, a substituent containing a hydrophilic group or a substituent containing a biologically active moiety, provided that one of R_2 , R_3 , R_{12} , R_{13} , R_{14} , and R_{15} is a substituent containing a biologically active moiety and at least one of the remaining of R_3 , R_{12} , R_{13} , R_{14} and R_{15} is a substituent containing a hydrophilic group.

Another preferred class of conjugates according to the invention which is used in preferred embodiments of the diagnostic assay and assay elements according to the invention is represented by the formula

40

45

10

20

wherein R_{12} , R_{13} , R_{14} and R_{15} each independently is hydrogen, a substituent containing a hydrophilic group or a substituent containing a biologically avtive moiety, provided that one of R_2 , R_3 , R_{12} , R_{13} , R_{14} and R_{15} is a substituent containing a biologically active moiety and at least one of the remaining of R_3 , R_{12} , R_{13} , R_{14} and R_{15} is a substituent containing a hydrophilic group.

As stated previously, in the conjugates of the invention the biologically active moiety may be attached to the fluorescent dye moiety at various positions. In a preferred embodiment the biologically active moiety is included within substituent R_2 (Formula I). The biologically active moiety is preferably attached through a linkage which is a carboxamido piperazinyl derivative represented by the formula

$$-N \qquad N-C-(CH_2)_{\overline{3}} \qquad (IV)$$

35 Thus, in this preferred embodiment R₂ is represented by

0

45

50

30

$$\begin{array}{c}
0 \\
-S - N
\end{array}$$

$$N - C - (CH_2)_3 - BIO$$
(VI)

The attachment of the biologically active moiety to the linkage may be through any appropriate position on that moiety. For example, where the biologically active moiety is the ophylline it may be attached to the linkage through the 3- or 8- positions.

The labeled conjugates of the invention may be used in any of many applications such as, for example, in diagnostic assays such as competitive binding assays or immunoassays or in immune respons reactions employing labeled reagents. In a preferred embodiment, the conjugates are employed in drug

monitoring applications such as for theophylline, phenytoin or in hormone monitoring applications such as thyroxine (T4). The particular assays in which the conjugates of the invention find utility are well known, e.g., immunometric assays, competitive binding assays. and therefore extensive discussion of such assays is not required here. In such known diagnostic tests, or assays, a biological reaction or interaction results in the generation of a detectable signal. For instance, in a typical immunoassay carried out in a multilayer assay element, an analyte - containing sample comprising a body fluid such as serum or whole blood is applied to a surface of the element. The fluid is typically passed through a filtering medium to remove interfering species and/or cells and then diffuses to a layer containing the labeled biologically active conjugate to produce an immunological reaction or interaction in accordance with the particular system of the assay element and this reaction or interaction in turn generates a detectable signal which is a function of the analyte in the sample fluid. The signal so generated may in less sophisticated systems provide only a qualitative determination of the presence of analyte or, in more sophisticated systems, it may provide a semi-quantitative measurement of the analyte. In such a system, the biologically active conjugate containing the detectable signal - generating moiety may be a so-called label-protein conjugate, i.e., a protein such as an antigen, antibody or Fab fragment "labeled" with, or containing, the dye moiety.

A typical multilayer assay element according to the invention comprises a support layer, which may be transparent, carrying at least one reagent layer in which there is provided the labeled conjugate of the invention. Also, in the reagent layer(s) there is provided any other reagent which is necessary for the particular signal generating system being exploited for the particular assay. These elements may also include other layers to provide various functions which are known in the art such as, for example: a layer to receive the sample and provide an even distribution of the sample components to the underlying reagent layer(s); a light-blocking, or screen, layer to assist in the detection of the signal by separating the layer in which the signal generating species is located from other layers of the element thus preventing undesirable interference with the detection of the signal; etc.

The conjugates of the invention may also be used for fluorescent staining of cells. The cells may then be observed under a microscope, the presence of the fluorescent conjugate being indicative of the presence of a specific determinant site. Further, the conjugates may be used for the deletion, separation or other application in a fluorescent activated cell sorter.

The conjugates of the invention may be prepared by reactions which are known to those skilled in the art and these will be apparent from the specific examples provided below herein. Accordingly, extensive discussion of such processes is not required here.

The invention will now be described further in detail with respect to specific preferred embodiments by way of Examples, it being understood that these are intended to be illustrative only and the invention is not limited to the specific materials, compositions or processes described therein.

EXAMPLE I

25

35

45

50

55

A mixture of 1.9g (13.0 mmole) of 7-hydroxyquinoline and 100 mg Pt₂O in 75 ml of 95% ethanol was hydrogenated overnight. The mixture was suction filtered through diatomaceous earth and the filtrate evaporated under reduced pressure. The resulting reddish-brown oil slowly crystallized to afford a quantitative yield of

which was stored in the dark under nitrogen.

'H NMR (CDCl₃): δ 1.85 (quint., 2H, J=6 Hz), 2.65(t, 2H, J=6 Hz): 3.23 (t, 2H, 6 Hz): 4.65(b.s., 2H), 5.95 (d. 1H, J=3 Hz); 6.1(dd, 1H, J₁=90 Hz, J₂=3 Hz), 6.75 (d, 1H, J=9 Hz). M.S. (M⁺ 149).

The crude product (VII) (28 mmole) was combined neat in a dry flask with 4.16g (28 mmole) of phthalic anhydride and 1.9g (14 mmole) of fused ZnCl₂ and the resulting mixture was h ated in an oil bath at 150 °C under nitrogen for 2 hours. The solid mass was cooled to room temperature, pulverized and extracted with warm water and suction filtered. The resulting red solid was taken up in 250 ml of 18% ag. HCl, warmed on

a steam bath and several ml of conc. HCl added to reach solubility. The solution was slowly cooled to room temperature and then stored in a freezer for 2 days. The resulting crystalline product was suction filtered, washed with excess 1N HCl and dried under vacuum to give 2.65g (42% yield) of

10

15

as a purple powder. Titration with potassium-t-butoxide suggested that the product was a bis-HCl salt (M⁺ 20 410 free base).

Potassium-t-butoxide (1.56g, 13.9 mmole) was added in one portion to a stirred solution of 2.0g of the previous product (VIII) in 200 ml of dry DMF at 0°-5°C under nitrogen. A cyan color formed and subsequently dissipated. (Note: subsequently it was found to be more convenient to add the potassium-t-butoxide to a point where the cyan color persists and then add one more equivalent). Another 0.5g (4.5 mmole) of potassium-t-butoxide was added and the resulting cyan solution stirred for 45 minutes. To this solution there was added, dropwise over a 5 minute period, a solution of 2.2 ml (13.4 mmole) t-butylbromoacetate in 10 ml DMF and the resulting mixture warmed to room temperature. The solvent was then removed under high vacuum followed by taking up the residue in CH₂Cl₂, filtering through diatomaceous earth and evaporating the filtrate. The product was purified by chromatography on silica gel using 10% CH₃OH/CH₂Cl₂ as the eluent to give 2.0g (70% yield) of

35

40

45

as a magenta powder.

'H NMR (CDCl₃): 1.5 (s, 18H), 1.9 (m,4H), 2.55 (t, 4H, J=6 Hz), 3.4 (t, 4H, J=6 Hz], 3.4 (t, 4H, J=6 Hz), 3.9 (bs., 4H), 6.23 (s,2H), 6.34 (s, 2H), 7.1 - 7.2 (m, 1H], 7.5-7.75 (m, 2H), 8.0-8.2 (m, 1H).

50

A mixture of 370 mg (0.58 mmole) of (IX) and 193 mg (0.58 mmole) of

5

45

50

55

(which was made from the corresponding 8-carboxypropyl theophylline and piperazine via trimethylacetyl mixed anhydride coupling) in 10 ml DMF was cooled to -30 C. Diphenylphosphoryl azide (160 ul, 0.75 mmole) was added to the stirred solution under nitrogen and the bath allowed to warm slowly to room temperature. After 4 days the solution was poured into excess ethyl ether, the liquid decanted from the oil and the oil chromatographed on silica gel using 10-15% CH₃OH/CH₂Cl₂ to give 421 mg (61% yield) of

as the diphenylphosphate salt ($X = (PhO)_2 PO_2^-$); (m/e = 956 FAB+).

'H NMR (CDCl₃) was complex but consistent with the structure of the product.

A solution of 421 mg (0.35 mmole) of (XI) in 10 ml CH₂Cl₂ was cooled to about 5 °C and treated with 2 ml of trifluoroacetic acid. The resulting mixture was stirred under nitrogen for 30 minutes and then at room temperature overnight. The product was isolated by evaporation of the solvent and purified by chromatography on silica gel with 15% - 30% CH₃OH/10% AcOH/CH₂Cl₂ as the eluent to give the desired conjugate

where R' is -OH and R" is -O $^{\circ}$, as a hygroscopic solid in 63% yield: (M $^{+}$ 843); 'H NMR (CD $_{3}$ OD, CDCl $_{3}$) complex but consistent with counterion being internal carboxylate compensated); λ max (pH7 buffer) 568 nm (ϵ = 107,073).

 $C_{45}\,H_{46}\,N_8\,O_9$. (H2O)4 requires 59.07% C, 5.95% H and 12.25% N. Elemental analysis found 59.26% C, 5.71% H and 12.25% N.

EXAMPLE II

25

Conjugates XIII (XII where R' is -NH(CH₂)₂SO₃H) and R" is -NH(CH₂)₂SO₃Θ) and XIV (XII where R' = R" and each is -N(CH₃)CH₂(CHOH)₄ CH₂OH) were synthesized from XII via diphenylphosphoryl azide (DPPA) coupling with taurine and N-methyl-d-glucamine, respectively. Thus, 1 mmole of XII was combined with 3 mmoles of taurine (or 3 mmoles of N-methyl-d-glucamine) and 5 mmoles of triethylamine in the case of taurine (or 2 mmoles in the case of N-methyl-d-glucamine) in 1/2 ml of dry DMF. To the cooled solutions (-30 °C) there were added 3 mmoles DPPA in one portion, the bath slowly warmed to room temperature and stirred overnight. The products were isolated by evaporation under vacuum and purified by reverse phase chromatography.

Conjugate XIII exhibited λ max (pH7 buffer) 558 nm (ϵ = 74,900). Conjugate XIV exhibited λ max (pH7 buffer) 555 nm (ϵ = 60,300).

EXAMPLE III

To a flamed/N₂ cooled 100 ml round bottom flask equipped with an inlet for N₂ atmosphere there was added 2.0g (4.1 X 10⁻³ mole) of bis rhodamine HCl salt (VIII) along with 50 ml of dry DMSO (48 hours over 3A sieves). The flask was immersed in a water bath at ambient temperature and then 1.89g (1.68 X 10⁻² mole) of t-butoxide potassium salt were added in one portion with stirring. The red-violet colored reaction mixture was stirred for 2 hours at ambient temperature. Propane sultone (2.54g, 2.08 X 10⁻² mole) was added in one portion and the reaction mixture stirred for an additional 14 hours at ambient temperature. The crude reaction mixture was evaporated under high vacuum and the resulting solid was triturated with CH₃CN and then with ethyl ether prior to chromatographic purification using a reverse phase support, (C₁₈ silica prepared according to Journal of Organic Chemistry, 48, 1983, page 3589) and 30% CH₃OH/H₂O as the eluent to yield 2.3g (72% yield) of a magenta powder, tris(3-sulfopropyl) rhodamine, represented by the formula

50

The structure of the product was confirmed by a 300 MHz NMR spectrum.

15

50

55

The tris(3-sulfopropyl) rhodamine (2.3g) was stirred in 30 ml of aqueous 1N NaOH for 1 hour under nitrogen. To this solution there was added 30 ml of 1N HCl and the solvent then evaporated to dryness under high vacuum. The resulting solid was dissolved in a small volume of water and applied to a pad of C₁₈ silica. The salts were removed by washing with water. The desired bis (3-sulfopropyl) rhodamine carboxylic acid was removed from the pad by elution with CH₃OH, the solvent removed by evaporation under vacuum and the solid product triturated with CH₃CN followed by drying under vacuum to give 1.67g (86% yield) of bis (3-sulfopropyl) rhodamine carboxylic acid*, a purple powder represented by the formula

Bis (3-sulfopropyl) rhodamine carboxylic acid (50 mg, 7.67×10^{-5} mole) was added to a flamed/N₂ cooled round bottom flask together with 2-3 ml of dry DMF and stirred at ambient temperature under nitrogen. Triethylamine (64 μ l, 46.4 mg, 4.6 \times 10⁻⁴ mole) was added followed by pivaloyl chloride (28.2 μ l, 27.6 mg, 2.3 \times 10⁻⁴ mole) and the mixture stirred for 2 hours. The reaction was monitored by TLC using a piperidine quench on aliquots. Piperazine (66 mg, 7.67 \times 10⁻⁴ mole) was added to the formed mixed anhydride and the reaction mixture stirred for an additional 14 hours. The solvent was then removed under high vacuum and the crude mixture was purified on a silica gel column using C₁₈ silica with 30% CH₃OH/H₂O as the eluent to give bis (3-sulfopropyl) rhodamine piperazinamide, represented by the formula

^{*}The compound is hygroscopic and should be stored under nitrogen.

The structure of the product was confirmed by NMR spectral data (300 MHz). Quantitative analysis was consistent with a hexahydrate. The product deliquesces readily and should be stored under high vacuum or nitrogen.

The mixed anhydride of 3-(carboxylpropyl) theophylline (which was prepared following the procedure described in Jour. of Org. Chem., 45 (9) 1980, page 1711) was formed in a flamed/N₂ cooled round bottom flask under nitrogen by addition of pivaloyl chloride (14.5 μl, 14.5 mg, 1.20 X 10⁻⁴ mole) and triethylamine (16.7 μl, 12.1 mg, 1.20 X 10⁻⁴ mole) to 3-(carboxypropyl) theophylline (28.6 mg, 1.20 X 10⁻⁴ mole) in 6 ml of dry DMF at 0 °C. The mixture was stirred for 1 1/2 hours at 0 °C and then transferred to a separate flamed/N₂ cooled round bottom flask containing bis (3-sulfopropyl) rhodamine piperazinamide (43.4 mg, 6.01 X 10⁻⁵ mole) under nitrogen. The reaction mixture was stirred for 12 hours at ambient temperature. Thin layer chromatography showed an incomplete reaction so 2 more equivalents of the mixed anhydride were added to the reaction mixture. The solvent was removed under high vacuum and the crude product. was purified by chromatography in accordance with the procedure previously described using 40% CH₃OH/H₂O as the eluent to give 40 mg (68.7% yield) of the rhodamine/3-(carboxypropyl) theophylline conjugate represented by the formula

where BIO is

15

35

40

50

55

The structure of the product was confirmed by a 300 MHz NMR spectrum, m/e = 958 FAB *.

EXAMPLE IV

L - thyroxine sodium salt, from Cal Biochem, (200 mg, 0.25 mmole) was added to a stirred solution of acetic anhydride (24 μ I, 0.25 mmole) in 2 ml DMF at room temperature under nitrogen. Triethylamine (35 μ I, 0.25 mmole) was then added and the resulting mixture stirred overnight. The solvent was removed under high vacuum, the residue taken up in 2 ml CH₃OH and then 1N HCl added dropwise, with stirring, to form a white precipitate. The supernatent liquid was decanted and the solid dried under vacuum to give 180 mg (88% yield) of

10

15

(XIX)

20

'H NMR (CD₃OD - CDCl₃): 2.02 (s, 3H), 2.8 - 3.2 (m, 2H), 4.70 (b.t., 1H, J=7 Hz), 7.1 (s, 2H), 7.7 (s, 2H); M.S. (M⁺ 819).

Trimethylacetyl chloride (30 μ l, 0.24 mmole) was added dropwise to a stirred solution of 180 mg (0.22 mmole) of XIX and triethylamine (34 μ l, 0.24 mmole) in 2 ml DMF at 0 $^{\circ}$ - 5 $^{\circ}$ C. The resulting mixture was stirred under nitrogen for 1 hour and then added dropwise to a cold solution of 95 mg (1.10 mmole) piperazine in 1 ml DMF. The mixture was warmed to room temperature, stirred for 2 hours and then evaporated to dryness. The residue was triturated with CH₃OH, filtered, triturated with H₂O, filtered and dried to give 41 mg (21% yield) of

30

35

40

M.S. (M+ 887).

A mixture of IX (41 mg, 0.064 mmole) and 57 mg of XX (0.064 mmole) in 1 ml DMF, at -30 °C under nitrogen, was treated in one portion with 18 μl (0.083 mmole) of diphenylphosphoryl azide. The resultant mixture was allowed to warm to room temperature and stirred overnight. The volatile matter was removed under high vacuum and the residue chromatographed on silica gel with 10% AcOH/20% CH₃OH/CH₂Cl₂ to give 12 mg (13% yield) of

50

(XXI)

conjugate XXII (XXI) where R' = R" and each is -OC(CH₃)₃ and X is diphenylphosphate).

Conjugate XXIII (XXI where R' is -OH and R" is O°) was made from XXII following the same procedure used for XII

EXAMPLE V

20

35

1-(carboxyethyl)phenobarbital (50 mg, 1.64 X 10-4 mole) was dissolved in 2 ml of dry DMF with stirring under N_2 . The solution was cooled in an ice water bath and triethylamine (25 μ l, 1.75 X 10⁻⁴ mole) added in one portion. Pivaloyl chloride (22 μ l, 1.75 X 10⁻⁴ mole) was added to the mixture in one portion. Within several minutes a precipitate was observed and the mixture was stirred at 5 °C for 1 hour. The mixture was then added slowly, dropwise, to a solution of piperazine (70.6 mg, 8.20 X 10⁻⁴ mole) in 5 ml of dry DMF with stirring under N_2 . The reaction mixture was stirred with cooling in an ice water bath for 1 hour and then at room temperature for 1 hour followed by concentration of the solvent under high vacuum at room temperature. The semi-solid residue was charged onto a silica gel column and eluted with 10% CH₃OH/1% NH₄OH/CH₂Cl₂. The desired product fractions were concentrated under vacuum to give 45.1 mg (74% yield) of

HN
$$N(CH_2)_2$$
 $C-N$
 C_2H_5

(XXIV)

Triethylamine (19 μ I, 1.33 X 10⁻⁴ mole) was added in one portion to a cooled solution (~ 3°C) of 78 mg (1.21 X 10⁻⁴ mole) of IX in 2 ml of dry DMF. Pivaloyl chloride (17 μ I, 1.33 X 10⁻⁴ mole) was added to the solution in one portion with cooling. A precipitate was observed almost immediately and the mixture was stirred for 1 hour at reduced temperature. The mixture was then added quickly, dropwise, to a solution of 45 mg (1.21 X 10⁻⁴ mole) of XXIV and triethylamine (19 μ I, 1.33 X 10⁻⁴ mole) in 2 ml of dry DMF. The resulting solution was stirred for 1 hour at reduced temperature and 1 hour at room temperature. The solution was then concentrated to a solid under high vacuum. The solid was purified on a silica gel column by eluting with 1:1 ethyl acetate/acetone to give the unreacted dy (24 mg, 31%). Elution with 10% CH₃OH/CH₂Cl₂ gave 65.3 mg (54.3%) of

XXVI (XXV where R' = R" and each is $-OC(CH_3)_3$) and X is chloride).

Compound XXVI (65.3 mg, 6.57 X 10⁻⁵ mole) was stirred in 1.5 ml of trifluoroacetic acid at room temperature under nitrogen for 1 1/2 hours and the solvent then concentrated under vacuum. The residue was charged onto reverse phase silica gel and eluted with 40% CH₃OH/H₂Oand then with 60% CH₃OH/H₂O to give numerous fractions with mixtures of products. The fractions exhibiting R_I = 0.79 with reverse phase silica gel were concentrated under vacuum and dried to give 6.1 mg of XXVII (XXV where R' is -OH and R" is -O^e). Subsequent syntheses indicated that elution from normal phase silica gel with 10% CH₃OH/10% The state of the state of the HOAc/CH₂Cl₂ provides better recovery. W. S. J. P. 18 82 1

EXAMPLE VI

20

30

40

45

50

A flask containing 40 ml of glacial acetic acid was cooled in a cold water bath and 1.9g (0.03 mole) of sodium cyanoborohydride was added portionwise with stirring and constant cooling. After H2 evolution had ceased 1.0g (0.0075 mole) of 4-hydroxyindole was added and the resulting solution was stirred at room temperature for 3 hours.

The reaction mixture was then slowly poured into a stirred solution of 50g sodium carbonate in 400 ml H₂O. The crude product was extracted out with ethyl acetate. The solvents were removed on the rotary evaporator and the residue was chromatographed on silica gel (10% EtOAc/CH₂Cl₂) to yield 723 mg (71%) of

4-hydroxyindoline (XXVIII) as a white powder.

¹H NMR (CDCl₃): δ 2.83 (t, 2H), 3.43 (t, 2H), 5.13 (b.s., 1H), 5.9 - 6.1 (m, 2H), 6.77 (t, 1H), 8.80 (s, 1H) m/e = 135.

A suspension of 1.6g (12.1 mmole) of mono-methylsuccinate in 20 ml of dry toluene was stirred at room temperature and 140 mg (3.7 mmole) of sodium borohydride was added. No evolution of gas was observed. The suspension was treated with 100 mg (0.74 mmole) of XXVIII and the temperature of the reaction mixture was slowly raised to 60 °C, during which time gas evolution started and total solution was achieved.

After 2 hours at 60°, the reaction solution was cooled to room temperature, then guenched into 200 ml saline, and the product was extracted out with EtOAC. The solvents wer removed on the rotary evaporator to give a crude product which was chromatographed on silica gel (5% EtOAC/CH2Cl2). The product containing fractions wer combined and evaporated, leaving a white solid. This solid was dissolved in CH₂Cl₂ and washed with 5% sodium bicarbonate solution. The layer was dried over sodium sulfate and then evaporated to yield 35 mg (20%) of XXIX as a pale oil.

15

¹H NMR (CDCl₃): δ 1.89 (q, 2H), 2.39 (t, 2H), 2.75 - 3.2 (m, 4H), 3.37 (t, 2H), 3.63 (s, 3H) 5.95 - 6.15 (m, 2H), 6.88 (t, 1H) m/e = 236 (FAB⁺).

Compound XXIX (32 mg, 0.14 mmole) was combined neat in a dry flask with phthalic anhydride (20.1 mg, 0.14 mmole) and fused ZnCl₂ (9 mg, 0.07 mmole) and the resulting mixture was heated in an oil bath at 150 °C under nitrogen for two hours. The resulting red mass was cooled to room temperature, then chromatographed on silica gel, using 3% MeOH - 5% MeOH/CH₂Cl₂ as eluant. The product containing fractions were combined and evaporated to yield 18 mg (45%) of XXX as a purple solid.

25
 $^{\text{CH}_3\text{O}_2\text{C}-(\text{CH}_2)_3-N}$ $^{\text{O}}$ $^{\text{N}-(\text{CH}_2)_3\text{CO}_2\text{CH}_3}$ $^{\text{30}}$ $^{\text{CO}_2}$

¹H NMR (CDCl₃ + CD₃OD): δ 1.97 (m, 4H), 2.40 (t, 4H), 3.3 - 3.7 (m, 8H), 3.63 (s, 6H), 4.03 (t, 4H), 6.57 (d, 2H, J = 9 Hz), 7.02 (d, 2H, J = 9 Hz), 7.3 (m. 1H), 7.75 (m, 2H), 8.3 (m, 1H). m/e = 583

A mixture of 18 mg (0.031 mmole) of XXX and 10.3 mg (0.031 mmole) of compound X in 1.5 ml DMF was cooled to -30 $^{\circ}$ C. Diphenylphosphoryl azide (10 mg, 0.036 mmole) was added to the stirred solution under nitrogen and the reaction mixture was slowly allowed to warn to ambient temperature, then stirred for 60 hours. The solvents were removed under high vacuum and the purple residue was chromatographed on silica gel (5% MeOH - 15% MeOH/CH₂Cl₂) to yield 32 mg (91%) of XXXI as a purple solid.

¹H NMR (CDCl₃ + CD₃OD) was complex but consistent with the structure of the product.

A solution of 30 mg (0.026 mmole) of XXXI in 2 ml ethanol was treated dropwise with 1.0 ml of 0.1 N NaOH solution and the resulting solution was heated at 30 °C under nitrogen until TLC showed complete hydrolysis (about 2 hours). The excess solvents were removed under high vacuum. The residue was stirred with 5% HOAc solution, and the excess solvents were again removed under vacuum. The crude product was purified by chromatography on silica gel with 10% - 20% MeOH/5% HOAc/CH₂Cl₂ as the eluant to yield 14 mg (62%) of the conjugate XXXII as a purple powder.

(XXXXI)

 $m/e = 872 (FAB^{+}).$

Visible spectrum: λ max (pH 7 buffer): 600 nm, ϵ = 58000.

¹H NMR (CD₃OD + CDCl₃) was complex but consistent with the counterion being internal carboxylate sompensated.

EXAMPLE VII

15

To further illustrate the utility of the biological fluorescent conjugates of the invention a thyroxine (T4) immunoassay was carried out. The following reagents were used in the assay:

Reagent A solution: 50 mmoles HEPES buffer (pH 7.2); 10 mmoles Na₂ EDTA; 0.02% 8-anilino-naphthalene sulfonate (ANS); 0.1% bovine serum albumin (BSA); and 0.05% NaN₃.

Reagent B solution: reagent A and 5% polyethylene glycol (PEG; NW 6000).

Calibrator solutions were made having 1, 25, 50, 100, 200 and 400 ng T4/ μ I of Reagent, A, respectively. The calibrator solutions (200 μ I) were each combined with 200 μ I of mouse monoclonal anti-T4 antibody (Behringwerke AG 49/7), 0.15 mmole Ig G, in Reagent A and 200 μ I of a solution of conjugate XXIII (prepared according to Exampl IV), 0.17 mmole, in Reagent A. The mixtur was incubated for 60 minutes at room temperatur. Subsequently, 200 μ I of 2% mouse normal serum in Reagent B and 200 μ I

of goat anti-mous Ig G antibody (1/16 dilution of Calbiochem Cat. No. 401210) in Reagent B wer added and the mixture further incubated for 15 minutes at room temperature.

The mixtures wer then centrifuged for 15 minutes at 3000g and the supernatant liquid decanted. The fluorescence of the free conjugate in the supernatant liquid was measured on a Perkin Elmer MPF 44 Fluorophotometer with excitation at 565 nm and the emission at 590 nm measured. Table I shows the results obtained:

TABLE I

	Thyroxin	Free Conjugate
	(ng/ul)	(%)
15	1	19.0
	25	25.4
	50	39.8
	100	71.8
20	200	84.2
	400	87.0

The standard curve exhibited good dose response in the 70-110 ng T4/ml range of interest.

Although the invention has been described with respect to specific preferred embodiments, the rings of the rhodamine label moiety of the conjugates can be for example appropriately substituted further such as with the solubilizing groups disclosed.

0 Claims

25

35

10

Claims for the following Contracting States: AT, BE, CH, DE, FR, GB, IT, LI, LU, NL, SE

 A fluorescent diagnostic assay comprising combining in an assay medium: a sample suspected of containing an analyte; a binding partner for said analyte; and a conjugate of said analyte and a fluorescent compound, said conjugate represented by the formula

$$\begin{array}{c|c} & -xm & -wp & \\ & & &$$

50

55

45

wherein W, X, Y and Z each independently is

-(-CH₂)_a(CHR₄) (CH₂-)-_b;

R, R₁, R₃ and R₄ each independently is hydrogen, a substituent containing a hydrophilic group or a substituent containing a biologically active moiety:

R₂ is -CO₂⁻, SO₃⁻ or a substituent containing a biologically active moiety;

provided that one of R, R_1 , R_2 , R_3 and R_4 is a substituent containing a biologically active moiety and at least one of the remaining of R, R_1 , R_3 and R_4 is a substituent containing a hydrophilic group;

a and b each independently is an integer of from 0 to 4, provided that the sum of a + b is an integer of from 1 to 4;

m and n each is 0 or 1 provided that one of m and n is 0 and the other is 1;

p and q each is 0 or 1 provided that one of p and q is 0 and the other is 1;

5

10

15

20

25

30

35

55

A is a counterion or the counterions to balance the overall charges on the conjugate moiety; and X is 0 or 1;

separating bound conjugate from free conjugate; measuring the level of the fluorescent signal of said bound or free conjugate; and

relating said level of said signal to the amount of said analyte in said sample.

2. The assay as defined in claim 1 wherein said conjugate is represented by the formula

$$\begin{array}{c|c}
R_{12} & R_{13} \\
\hline
R_{14} & R_{15}
\end{array}$$

$$\begin{array}{c}
R_{13} \\
R_{15}
\end{array}$$

$$\begin{array}{c}
R_{15} \\
R_{3}
\end{array}$$

wherein R_{12} , R_{13} , R_{14} and R_{15} each independently is hydrogen, a substituent containing a hydrophilic group or a substituent containing a biologically active moiety, provided that one of R_2 , R_3 , R_{12} , R_{13} , R_{14} , and R_{15} is a substituent containing a biologically active moiety and at least one of the remaining of R_3 , R_{12} , R_{13} , R_{14} and R_{15} is a substituent containing a hydrophilic group.

3. The assay as defined in claim 1 wherein said conjugate is represented by the formula

$$R_{12} \longrightarrow R_{13} \longrightarrow R$$

wherein R₁₂, R₁₃, R₁₄ and R₁₅ each independently is hydrogen, a substituent containing a hydrophilic group or a substituent containing a biologically active moiety, provided that one of R₂, R₃, R₁₂, R₁₃,

EP 0 285 179 B1

R₁₄ and R₁₅ is a substituent containing a biologically active moiety and at least one of the remaining of R₃, R₁₂, R₁₃, R₁₄ and R₁₅ is a substituent containing a hydrophilic group.

- 4. The assay as defined in claims 2 or 3 wherein R₂ is a substituent containing a biologically active moiety.
 - The assay as defined in any of claims 2 or 3 wherein R₁₂ and/or R₁₃ is a substituent containing a hydrophilic group
- 70 6. The assay as defined in claim 5 wherein said hydrophilic group is a carboxylic acid, a polyether, a polyalcohol, an amine, a sulfonic acid, a phosphonic acid, a phosphonic ester, a phosphate, a phosphate ester, a phosphinic acid, a boronic acid or a borinic acid.
- 7. A diagnostic assay element adapted to receive a sample of a biological fluid and to provide a detectable signal as a function of an analyte which may be present in said fluid, said assay element containing a fluorescent conjugate represented by the formula

wherein W, X, Y and Z each independently is

-(-CH₂)_a(CHR₄) (CH₂-)-_b;

R, R₁, R₃ and R₄ each independently is hydrogen, a substituent including a hydrophilic group or a substituent containing a biologically active moiety selected from an antigen, an antibody, a hapten, an Fab fragment and a DNA probe;

R₂ is -CO₂⁻, SO₃⁻ or a substituent containing said biologically active moiety;

provided that one of R, R_1 , R_2 , R_3 and R_4 is a substituent containing said biologically active moiety and at least one of the remaining of R, R_1 , R_3 and R_4 is a substituent containing a hydrophilic group;

a and b each independently is an integer of from 0 to 4, provided that the sum of a + b is an integer of from 1 to 4;

m and n each is 0 or 1 provided that one of m and n is 0 and the other is 1;

p and q each is 0 or 1 provided that one of p and q is 0 and the other is 1;

- A is a counterion or the counterions to balance the overall charges on the conjugate moiety; and X is 0 or 1.
 - 8. The diagnostic element as defined in claim 7 which contains a support carrying at least one reagent layer.
 - 9. The diagnostic element as defined in claim 8 which contains the binding partner of said biologically active moiety present in said conjugate.

5**5**

50

35

10. The assay element as defined in claim 7 wherein said conjugate is represented by the formula

5

10

15

20

25

50

55

wherein R_{12} , R_{13} , R_{14} and R_{15} each independently is hydrogen, a substituent containing a hydrophilic group or a substituent containing said biologically active moiety, provided that one of R_2 , R_3 , R_{12} , R_{13} , R_{14} and R_{15} is a substituent containing said biologically active moiety and at least one of the remaining of R_3 , R_{12} , R_{13} , R_{14} and R_{15} is a substituent including a hydrophilic group.

11. The assay element as defined in claim 7, wherein said conjugate is represented by the formula

30
$$R_{12}$$

$$R_{14}$$

$$R_{15}$$

$$R_{16}$$

$$R_{18}$$

$$R_{19}$$

$$R_{19}$$

$$R_{10}$$

wherein R₁₂, R₁₃, R₁₄ and R₁₅ each independently is hydrogen, a substituent containing a hydrophilic group or a substituent containing said biologically active moiety, provided that one of R₂, R₃, R₁₂, R₁₃, R₁₄ and R₁₅ is a substituent containing said biologically active moiety and at least one of the remaining of R₃, R₁₂, R₁₃, R₁₄ and R₁₅ is a substituent containing a hydrophilic group.

- 12. The assay element as defined in claims 10 or 11 wherein R₂ is a substituent containing said biologically active moiety.
- 13. The assay element as defined in any of claims 10 or 11 wherein R₁₂ and/or R₁₃ is a substituent containing a hydrophilic group.

- 14. The assay element as defined in claim 13 wherein said hydrophilic group is a carboxylic acid, a polyether, a polyalcohol, an amine, a sulfonic acid, a phosphonic acid, a phosphonic ester, a phosphate, a phosphate ester, a phosphinic acid, a boronic acid or a borinic acid.
- 5 15. A fluorescent conjugate represented by the formula

20

15

wherein W, X, Y and Z each independently is

-(-CH₂)_a(CHR₄) (CH₂-)-_b;

 R_1 , R_3 and R_4 each independently is hydrogen, a substituent containing a hydrophilic group or a substituent containing a biologically active moiety selected from an antigen, an antibody, a hapten, a Fab fragment and a DNA probe;

R₂ is -CO₂⁻, SO₃⁻ or a substituent containing said biologically active moiety;

provided that one of R, R_1 , R_2 , R_3 and R_4 is a substituent containing said biologically active moiety and at least one of the remaining of R, R_1 , R_3 and R_4 is a substituent containing a hydrophilic group;

a and b each independently is an integer of from 0 to 4, provided that the sum of a + b is an integer of from 1 to 4;

m and n each is 0 or 1 provided that one of m and n is 0 and the other is 1;

p and q each is 0 or 1 provided that one of p and q is 0 and the other is 1;

A is a counterion or the counterions to balance the overall charges on the conjugate moiety; and X is 0 or 1.

35

30

16. The fluorescent conjugate as defined in claim 15 which is represented by the formula

45 R_{14} R_{15} R_{15} R_{15}

(A) ;

5**5**

wherein R₁₂, R₁₃, R₁₄ and R₁₅ each independently is hydrogen, a substituent containing a hydrophilic group or a substituent containing said biologically active moiety, provided that one of R₂, R₃, R₁₂, R₁₃,

R₁₄ and R₁₅ is a substituent containing said biologically active moiety and at least one of the remaining of R₃, R₁₂, R₁₃, R₁₄ and R₁₅ is a substituent containing a hydrophilic group.

17. The fluorescent conjugate as defined in claim 15 which is represented by the formula

- wherein R₁₂, R₁₃, R₁₄ and R₁₅ each independently is hydrogen, a substituent containing a hydrophilic group or a substituent containing said biologically active moiety, provided that one of R₂, R₃, R₁₂, R₁₃, R₁₄ and R₁₅ is a substituent containing said biologically active moiety and at least one of the remaining of R₃, R₁₂, R₁₃, R₁₄ and R₁₅ is a substituent containing a hydrophilic group.
- 18. The fluorescent conjugate as defined in any of claims 16 or 17 wherein R is a substituent containing said biologically active moiety.
 - 19. The use of the fluorescent conjugate as defined in any of claims 15 to 18, for fluorescent staining of cells, or for the detection or separation in a fluorescent activated cell sorter.

Claims for the following Contracting States: ES, GR

35

40

 A fluorescent diagnostic assay comprising combining in an assay medium: a sample suspected of containing an analyte; a binding partner for said analyte; and a conjugate of said analyte and a fluorescent compound, said conjugate represented by the formula

wherein W, X, Y and Z each independently is

EP 0 285 179 B1

(CH₂)_a(CHR₄) (CH₂)_b;

40

45

50

55

R, R₁, R₂ and R₄ each independently is hydrogen, a substituent containing a hydrophilic group or a substituent containing a biologically activ moiety;

R₂ is -CO₂⁻, SO₃⁻ or a substituent containing a biologically active moiety;

provided that one of R, R₁, R₂, R₃ and R₄ is a substituent containing a biologically active moiety and at least one of the remaining of R, R₁, R₃ and R₄ is a substituent containing a hydrophilic group; a and b each independently is an integer of from 0 to 4, provided that the sum of a + b is an integer of

from 1 to 4;

m and n each is 0 or 1 provided that one of m and n is 0 and the other is 1; p and q each is 0 or 1 provided that one of p and q is 0 and the other is 1;

A is a counterion or the counterions to balance the overall charges on the conjugate moiety; and X is 0 or 1:

separating bound conjugate from free conjugate; measuring the level of the fluorescent signal of said bound or free conjugate; and

- relating said level of said signal to the amount of said analyte in said sample.
 - 2. The assay as defined in claim 1 wherein said conjugate is represented by the formula

$$\begin{array}{c}
R_{12} \\
R_{14}
\end{array}$$

$$\begin{array}{c}
R_{13} \\
R_{15}
\end{array}$$

$$\begin{array}{c}
R_{13} \\
R_{15}
\end{array}$$

$$\begin{array}{c}
R_{15} \\
R_{3}
\end{array}$$

$$\begin{array}{c}
R_{13} \\
R_{15}
\end{array}$$

wherein R_{12} , R_{13} , R_{14} and R_{15} each independently is hydrogen, a substituent containing a hydrophilic group or a substituent containing a biologically active moiety, provided that one of R_2 , R_3 , R_{12} , R_{13} , R_{14} , and R_{15} is a substituent containing a biologically active moiety and at least one of the remaining of R_3 , R_{12} , R_{13} , R_{14} and R_{15} is a substituent containing a hydrophilic group.

3. The assay as defined in claim 1 wherein said conjugate is represented by the formula

5

10

15

20

25

40

wherein R_{12} , R_{13} , R_{14} and R_{15} each independently is hydrogen, a substituent containing a hydrophilic group or a substituent containing a biologically active moiety, provided that one of R_2 , R_3 , R_{12} , R_{13} , R_{14} and R_{15} is a substituent containing a biologically active moiety and at least one of the remaining of R_3 , R_{12} , R_{13} , R_{14} and R_{15} is a substituent containing a hydrophilic group.

- 4. The assay as defined in claims 2 or 3 wherein R₂ is a substituent containing a biologically active moiety.
- 5. The assay as defined in any of claims 2 or 3 wherein R and/or R₁₃ is a substituent containing a hydrophilic group
 - 6. The assay as defined in claim 5 wherein said hydrophilic group is a carboxylic acid, a polyether, a polyalcohol, an amine, a sulfonic acid, a phosphonic acid, a phosphonic ester, a phosphate, a phosphate ester, a phosphinic acid, a boronic acid or a borinic acid.
 - 7. A diagnostic assay element adapted to receive a sample of a biological fluid and to provide a detectable signal as a function of an analyte which may be present in said fluid, said assay element containing a fluorescent conjugate represented by the formula

wherein W, X, Y and Z each independently is -(-CH₂)_a(CHR₄) (CH₂-)-_b;

R, R₁, R₃ and R₄ each independently is hydrogen, a substituent including a hydrophilic group or a substituent containing a biologically active moiety selected from an antigen, an antibody, a hapten, an

EP 0 285 179 B1

Fab fragment and a DNA probe;

R₂ is -CO₂⁻, SO₃⁻ or a substituent containing said biologically active moiety;

provided that one of R, R_1 , R_2 , R_3 and R_4 is a substituent containing said biologically active moiety and at least one of the remaining of R, R_1 , R_3 and R_4 is a substituent containing a hydrophilic group;

a and b each independently is an integer of from 0 to 4, provided that the sum of a + b is an integer of from 1 to 4;

m and n each is 0 or 1 provided that one of m and n is 0 and the other is 1;

p and q each is 0 or 1 provided that one of p and q is 0 and the other is 1;

A is a counterion or the counterions to balance the overall charges on the conjugate moiety; and X is 0 or 1.

- The diagnostic element as defined in claim 7 which contains a support carrying at least one reagent layer.
- The diagnostic element as defined in claim 8 which contains the binding partner of said biologically active moiety present in said conjugate.
 - 10. The assay element as defined in claim 7 wherein said conjugate is represented by the formula

25

30

10

35

40

wherein R_{12} , R_{13} , R_{14} and R_{15} each independently is hydrogen, a substituent containing a hydrophilic group or a substituent containing said biologically active moiety, provided that one of R_2 , R_3 , R_{12} , R_{13} , R_{14} and R_{15} is a substituent containing said biologically active moiety and at least one of the remaining of R_3 , R_{12} , R_{13} , R_{14} and R_{15} is a substituent including a hydrophilic group.

45

50

11. The assay element as defined in claim 7, wherein said conjugate is represented by the formula

wherein R_{12} , R_{13} , R_{14} and R_{15} each independently is hydrogen, a substituent containing a hydrophilic group or a substituent containing said biologically active moiety, provided that one of R_2 , R_3 , R_{12} , R_{13} , R_{14} and R_{15} is a substituent containing said biologically active moiety and at least one of the remaining of R_3 , R_{12} , R_{13} , R_{14} and R_{15} is a substituent containing a hydrophilic group.

- 12. The assay element as defined in claims 10 or 11 wherein R_2 is a substituent containing said biologically active moiety.
- 13. The assay element as defined in any of claims 10 or 11 wherein R₁₂ and/or R₁₃ is a substituent containing a hydrophilic group.
 - 14. The assay element as defined in claim 13 wherein said hydrophilic group is a carboxylic acid, a polyether, a polyalcohol, an amine, a sulfonic acid, a phosphonic acid, a phosphonic ester, a phosphate, a phosphate ester, a phosphinic acid, a boronic acid or a borinic acid.
 - 15. The use of the fluorescent conjugate as defined in any of claims 1 to 3, for fluorescent staining of cells, or for the detection or separation in a fluorescent activated cell sorter.

Patentansprüche

5

10

15

20

25

45

50

55

Patentansprüche für folgende Vertragsstaaten : AT, BE, CH, DE, FR, GB, IT, LI, LU, NL, SE

 Diagnostischer Fluoreszenz-Assay, wonach in einem Assay-Medium kombiniert werden: eine Probe, die vermutlich einen Analyten enthält; einen Bindungspartner für den Analyten; und ein Konjugat des Analyten und einer fluoreszierenden Verbindung, wobei das Konjugat durch die Formel

dargestellt ist, worin W, X, Y und Z jeweils unabhängig voneinander

R, R₁, R₃ und R₄ jeweils unabhängig voneinander Wasserstoff, einen Substituenten mit einer hydrophilen Gruppe, oder einen Substituenten mit einer biologisch aktiven Gruppierung;

 R_2 die Gruppen $-CO_2^-$, SO_3^- oder einen Substituenten mit einer biologisch aktiven Gruppierung, darstellen;

vorausgesetzt, daß ein R, R₁, R₂, R₃ und R₄ einen Substituenten mit einer biologisch aktiven Gruppierung und mindestens ein verbleibendes R, R₁, R₂, R₃ und R₄ einen Substituenten mit einer hydrophilen Gruppe darstellen;

a und b unabhängig voneinander eine ganze Zahl von 0 bis 4 bedeuten, vorausgesetzt, daß die Summe von a + b eine ganze Zahl von 1 bis 4 darstellt;

m und n jeweils 0 oder 1 bedeuten, vorausgesetzt, daß ein m und n = 0 und das andere = 1 bedeutet; p und q jeweils 0 oder 1 bedeuten, vorausgesetzt, daß ein p und q = 0 und das andere = 1 bedeutet; A ein Gegenion oder die Gegenionen zum Ausgleich der Gesamtladungen auf der Konjugatgruppierung darstellt;

X = 0 oder 1;

10

15

25

30

35

40

45

50

55

worauf das gebundene Konjugat vom freien Konjugat getrennt wird; der Wert des Fluoreszenzsignals des gebundenen oder freien Konjugats gemessen wird; und der Wert des Signals mit der Menge des Analyten in der Probe in Beziehung gesetzt wird.

2. Assay nach Anspruch 1, worin das Konjugat durch die Formel

dargestellt ist, worin R_{12} , R_{13} , R_{14} und R_{15} unabhängig voneinander Wasserstoff, einen Substituenten mit einer hydrophilen Gruppe oder einen Substituenten mit einer biologisch aktiven Gruppierung darstellen, vorausgesetzt, daß ein R_2 , R_3 , R_{12} , R_{13} , R_{14} und R_{15} einen Substituenten mit einer biologisch aktiven Gruppierung und mindestens eine verbleibende Gruppe R_3 , R_{12} , R_{13} , R_{14} und R_{15} einen Substituenten mit einer hydrophilen Gruppe darstellt.

3. Assay nach Anspruch 1, worin das Konjugat durch die Formel

20

25

50

35
$$R_{12}$$
 R_{14} R_{15} R_{13} R_{14} R_{15} R_{13} R_{14} R_{15} R_{15}

dargestellt ist, worin R₁₂, R₁₃, R₁₄ und R₁₅ jeweils unabhängig voneinander Wasserstoff, einen Substituenten mit einer hydrophilen Gruppe oder einen Substituenten mit einer biologisch aktiven Gruppierung darstellen, vorausgesetzt, daß ein R₂, R₃, R₁₂, R₁₃, R₁₄ und R₁₅ einen Substituenten mit einer biologisch aktiven Gruppierung und mindestens eine verbleibende Gruppe R₃, R₁₂, R₁₃, R₁₄ und R₁₅ einen Substituenten mit einer hydrophilen Gruppe darstellt.

4. Assay nach Anspruch 2 oder 3, worin R₂ einen Substituenten mit einer biologisch aktiven Gruppierung darstellt.

- Assay nach einem der Ansprüche 2 oder 3, worin R₁₂und/oder R₁₃ einen Substituenten mit einer hydrophilen Gruppe darstellt.
- 6. Assay nach Anspruch 5, worin die hydrophile Gruppe eine Carbonsäure-, eine Polyether-, eine Polyalkohol-, eine Amino-, eine Sulfonsäure-, eine Phosphonsäure-, eine Phosphonester-, eine Phosphate-, eine Boronsäure- oder eine Borinsäuregruppe darstellt.
- 7. Diagnostisches Assay-Element zur Aufnahme einer Probe einer biologischen Flüssigkeit und zur 10 Erzeugung eines nachweisbaren Signals als Funktion eines Analyten, der in der Flüssigkeit vorhanden sein kann, wobei das Assay-Element ein fluoreszierendes Konjugat enthält, das durch die Formel

dargestellt ist, worin W, X, Y und Z jeweils unabhängig voneinander :

$$\begin{array}{cccc}
& \leftarrow \text{CH} & \text{CHR} & \text{CHR} & \text{CH} \\
& 2 & a & 4 & 2 & b
\end{array}$$

R, R₁, R₃ und R₄ jeweils unabhängig voneinander Wasserstoff, einen Substituenten mit einer hydrophilen Gruppe, oder einen Substituenten mit einer biologisch aktiven Gruppierung, ausgewählt aus einem Antigen, einem Antikörper, einem Hapten, einem Fab-Fragment und einer DNA-Sonde, bedeuten; R₂ die Gruppen -CO₂⁻, SO₃⁻ oder einen Substituenten mit der biologisch aktiven Gruppierung darstellen;

vorausgesetzt, daß ein R, R_1 , R_2 , R_3 und R_4 einen Substituenten mit der biologisch aktiven Gruppierung und mindestens ein verbleibendes R, R_1 , R_2 , R_3 und R_4 einen Substituenten mit einer hydrophilen Gruppe darstellen;

a und b unabhängig voneinander eine ganze Zahl von 0 bis 4 bedeuten, vorausgesetzt, daß die Summe von a + b eine ganze Zahl von 1 bis 4 darstellt;

m und n jeweils 0 oder 1 bedeuten, vorausgesetzt, daß ein m und n = 0 und das andere = 1 bedeutet; p und q jeweils 0 oder 1 bedeuten, vorausgesetzt, daß ein p und q = 0 und das andere = 1 bedeutet; A ein Gegenion oder die Gegenionen zum Ausgleich der Gesamtladungen auf der Konjugatgruppierung darstellt; und

X = 0 oder 1 ist.

5

35

40

45

- Diagnostisches Element nach Anspruch 7, enthaltend eine Unterlage, welche mindestens eine Reagensschicht trägt.
 - Diagnostisches Element nach Anspruch 8, enthaltend den Bindungspartner der im Konjugat vorliegenden biologisch aktiven Gruppierung.

10. Assay-Element nach Anspruch 7, worin das Konjugat durch die Formel

5

10

15

20

25

50

$$\begin{array}{c|c}
R_{12} & R_{13} \\
\hline
N & O & N \\
\hline
R_{14} & R_{15}
\end{array}$$

$$\begin{array}{c|c}
R_{13} & R_{15} \\
\hline
R_{3} & R_{15}
\end{array}$$

definiert ist, worin R_{12} , R_{13} , R_{14} und R_{15} unabhängig voneinander Wasserstoff, einen Substituenten mit einer hydrophilen Gruppe oder einen Substituenten mit der biologisch aktiven Gruppierung darstellen, vorausgesetzt, daß ein R_2 , R_3 , R_{12} , R_{13} , R_{14} und R_{15} einen Substituenten mit der biologisch aktiven Gruppierung und mindestens eine verbleibende Gruppe R_3 , R_{12} , R_{13} , R_{14} und R_{15} einen Substituenten mit einer hydrophilen Gruppe darstellt.

11. Assay-Element nach Anspruch 7, worin das Konjugat durch die Formel

30
$$R_{12} \longrightarrow R_{14} \longrightarrow R_{15} \longrightarrow R_{13}$$

$$R_{12} \longrightarrow R_{14} \longrightarrow R_{13} \longrightarrow R_{2} \longrightarrow R_{3}$$

$$R_{15} \longrightarrow R_{13} \longrightarrow R_{14} \longrightarrow R_{15} \longrightarrow$$

definiert ist, worin R₁₂, R₁₃, R₁₄ und R₁₅ jeweils unabhängig voneinander Wasserstoff, einen Substituenten mit einer hydrophilen Gruppe oder einen Substituenten mit der biologisch aktiven Gruppierung darstellen, vorausgesetzt, daß ein R₂, R₃, R₁₂, R₁₃, R₁₄ und R₁₅ einen Substituenten mit der biologisch aktiven Gruppierung und mindestens eine verbleibende Gruppe R₃, R₁₂, R₁₃, R₁₄ und R₁₅ einen Substituenten mit einer hydrophilen Gruppe darstellt.

- Assay-El ment nach Anspruch 10 oder 11, worin R₂ einen Substituenten mit der biologisch aktiven
 Gruppierung darstellt.
 - 13. Assay-Element nach einem der Ansprüche 10 oder 11, worin R₁₂ und/oder R₁₃ einen Substituenten mit einer hydrophilen Gruppe darstellen.

- 14. Assay-Element nach Anspruch 13, worin die hydrophile Gruppe eine Carbonsäure-, eine Polyether-, eine Polyalkohol-, eine Amino-, eine Sulfonsäure-, eine Phosphonsäure-, eine Phosphonsäure- oder eine Borinsäuregruppe darstellt.
- 15. Fluoreszierendes Konjugat, dargestellt durch die Formel

worin W, X, Y und Z jeweils unabhängig voneinander

$$+ CH_2$$
 (CHR₄) (CH₂ $+$ b)

R, R₁, R₃ und R₄ jeweils unabhängig voneinander Wasserstoff, einen Substituenten mit einer hydrophilen Gruppe, oder einen Substituenten mit einer biologisch aktiven Gruppierung aus der Gruppe bestehend aus einem Antigen, einem Antikörper, einem Hapten, einem Fab-Fragment und einer DNA-Sonde bedeuten;

R₂ die Gruppen -CO₂⁻, SO₃⁻ oder einen Substituenten mit der biologisch aktiven Gruppierung darstellen;

vorausgesetzt, daß ein R, R_1 , R_2 , R_3 und R_4 einen Substituenten mit der biologisch aktiven Gruppierung und mindestens ein verbleibendes R, R_1 , R_2 , R_3 und R_4 einen Substituenten mit einer hydrophilen Gruppe darstellen;

a und b unabhängig voneinander eine ganze Zahl von 0 bis 4 bedeuten, vorausgesetzt, daß die Summe von a + b eine ganze Zahl von 1 bis 4 darstellt;

m und n jeweils 0 oder 1 bedeuten, vorausgesetzt, daß ein m und n=0 und das andere = 1 bedeutet; p und q jeweils 0 oder 1 bedeuten, vorausgesetzt, daß ein p und q=0 und das andere = 1 bedeutet; A ein Gegenion oder die Gegenionen zum Ausgleich der Gesamtladungen auf der Konjugatgruppierung darstellt; und

X = 0 oder 1 ist.

50

25

30

35

40

16. Fluoreszierendes Konjugat nach Anspruch 15, dargestellt durch die Form 1

5

10

15

20

25

50

$$\begin{array}{c|c} & & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

worin R₁₂, R₁₃, R₁₄ und R₁₅ unabhängig voneinander Wasserstoff, einen Substituenten mit einer hydrophilen Gruppe oder einen Substituenten alt der biologisch aktiven Gruppierung darstellen, vorausgesetzt, daß ein R₂, R₃, R₁₂, R₁₃, R₁₄ und R₁₅ einen Substituenten mit der biologisch aktiven Gruppierung und mindestens eine verbleibende Gruppe R₃, R₁₂, R₁₃, R₁₄ und R₁₅ einen Substituenten mit einer hydrophilen Gruppe darstellt.

17. Fluoreszierendes Konjugat nach Anspruch 15, dargestellt durch die Formel

$$\begin{array}{c|c}
R_{14} & R_{15} \\
R_{12} & R_{13} \\
R_{3} & R_{15}
\end{array}$$

$$\begin{array}{c|c}
R_{14} & R_{15} \\
R_{13} & R_{13} \\
R_{2} & R_{3}
\end{array}$$

worin R₁₂, R₁₃, R₁₄ und R₁₅ jeweils unabhängig voneinander Wasserstoff, einen Substituenten mit einer hydrophilen Gruppe oder einen Substituenten mit der biologisch aktiven Gruppierung darstellen, vorausgesetzt, daß ein R₂, R₃, R₁₂, R₁₃, R₁₄ und R₁₅ einen Substituenten mit der biologisch aktiven Gruppierung und mindestens eine verbleibende Gruppe R₃, R₁₂, R₁₃, R₁₄ und R₁₅ einen Substituenten mit einer hydrophilen Gruppe darstellt.

- 18. Fluoreszierendes Konjugat nach einem der Ansprüche 16 oder 17, worin R₂ einen Substituenten mit der biologisch aktiven Gruppierung darstellt.
- 19. Verw ndung des fluoreszierenden Konjugats nach einem der Ansprüche 15 bis 18, zur Fluoreszenzfärbung von Zellen oder deren Nachweis oder Abtrennung in einer Sortiervorrichtung für fluoreszierende aktivierte Zellen.

Pat ntansprüche für folgende Vertragsstaaten: ES, GR

 Diagnostischer Fluoreszenz-Assay, wonach in einem Assay-Medium, kombiniert werden: eine Probe, die vermutlich einen Analyten enthält; einen Bindungspartner für den Analyten; und ein Konjugat des Analyten und einer fluoreszierenden Verbindung, wobei das Konjugat durch die Formel

dargestellt ist, worin W, X, Y und Z jeweils unabhängig voneinander

$$\sim$$
 CH₂ a (CHR₄) (CH₂ \rightarrow b)

R, R₁, R₃ und R₄ jeweils unabhängig voneinander Wasserstoff, einen Substituenten mit einer hydrophilen Gruppe, oder einen Substituenten mit einer biologisch aktiven Gruppierung;

R₂ die Gruppen -CO₂⁻, SO₃⁻ oder einen Substituenten mit einer biologisch aktiven Gruppierung, darstellen;

vorausgesetzt, daß ein R, R_1 , R_2 , R_3 und R_4 einen Substituenten mit einer biologisch aktiven Gruppierung und mindestens ein verbleibendes R, R_1 , R_2 , R_3 und R_4 einen Substituenten mit einer hydrophilen Gruppe darstellen;

a und b unabhängig voneinander eine ganze Zahl von 0 bis 4 bedeuten, vorausgesetzt, daß die Summe von a + b eine ganze Zahl von 1 bis 4 darstellt;

m und n jeweils 0 oder 1 bedeuten, vorausgesetzt, daß ein m und n=0 und das andere = 1 bedeutet; p und q jeweils 0 oder 1 bedeuten, vorausgesetzt, daß ein p und q=0 und das andere = 1 bedeutet; A ein Gegenion oder die Gegenionen zum Ausgleich der Gesamtladungen auf der Konjugatgruppierung darstellt;

X = 0 oder 1;

5

10

15

20

25

30

35

40

worauf das gebundene Konjugat vom freien Konjugat getrennt wird; der Wert des Fluoreszenzsignals des gebundenen oder freien Konjugats gemessen wird; und der Wert des Signals mit der Menge des Analyten in der Probe in Beziehung gesetzt wird.

2. Assay nach Anspruch 1, worin das Konjugat durch die Formel

5

10

15

20

25

50

$$\begin{array}{c|c}
R_{12} \\
N \\
R_{14}
\end{array}$$

$$\begin{array}{c}
R_{13} \\
R_{15}
\end{array}$$

$$\begin{array}{c}
R_{15} \\
R_{3}
\end{array}$$

$$\begin{array}{c}
R_{15} \\
R_{3}
\end{array}$$

dargestellt ist, worin R_{12} , R_{13} , R_{14} und R_{15} unabhängig voneinander Wasserstoff, einen Substituenten mit einer hydrophilen Gruppe oder einen Substituenten mit einer biologisch aktiven Gruppierung darstellen, vorausgesetzt, daß ein R_2 , R_3 , R_{12} , R_{13} , R_{14} und R_{15} einen Substituenten mit einer biologisch aktiven Gruppierung und mindestens eine verbleibende Gruppe R_3 , R_{12} , R_{13} , R_{14} und R_{15} einen Substituenten mit einer hydrophilen Gruppe darstellt.

3. Assay nach Anspruch 1, worin das Konjugat durch die Formel

$$R_{12}$$
 R_{14}
 R_{15}
 R_{13}
 R_{14}
 R_{15}
 R_{14}
 R_{15}
 R_{15}
 R_{16}
 R_{17}
 R_{18}
 R_{19}
 R

dargestellt ist, worin R_{12} , R_{13} , R_{14} und R_{15} jeweils unabhängig voneinander Wasserstoff, einen Substituenten mit einer hydrophilen Gruppe oder einen Substituenten mit einer biologisch aktiven Gruppierung darstellen, vorausgesetzt, daß ein R_2 , R_3 , R_{12} , R_{13} , R_{14} und R_{15} einen Substituenten mit einer biologisch aktiven Gruppierung und mindestens eine verbleibende Gruppe R_3 , R_{12} , R_{13} , R_{14} und R_{15} einen Substituenten mit einer hydrophilen Gruppe darstellt.

Assay nach Anspruch 2 oder 3, worin R₂ einen Substituenten mit einer biologisch aktiven Gruppierung darstellt.

- 5. Assay nach einem der Ansprüche 2 oder 3, worin R₁₂und/oder R₁₃ einen Substituenten mit einer hydrophilen Gruppe darstellt.
- 6. Assay nach Anspruch 5, worin die hydrophile Gruppe eine Carbonsäure-, eine Polyether-, eine Polyalkohol-, eine Amino-, eine Sulfonsäure-, eine Phosphonsäure-, eine Phosphonester-, eine Phosphate-, eine Phosphate-, eine Boronsäure- oder eine Borinsäuregruppe darstellt.
- 7. Diagnostisches Assay-Element zur Aufnahme einer Probe einer biologischen Flüssigkeit und zur Erzeugung eines nachweisbaren Signals als Funktion eines Analyten, der in der Flüssigkeit vorhanden sein kann, wobei das Assay-Element ein fluoreszierendes Konjugat enthält, das durch die Formel

dargestellt ist, worin W, X, Y und Z jeweils unabhängig voneinander

30
 \rightarrow CH₂) (CHR₄) (CH₂) 2 b

R, R₁, R₃ und R₄ jeweils unabhängig voneinander Wasserstoff, einen Substituenten mit einer hydrophilen Gruppe, oder einen Substituenten mit einer biologisch aktiven Gruppierung, ausgewählt aus einem Antigen, einem Antikörper, einem Hapten, einem Fab-Fragment und einer DNA-Sonde, bedeuten; R₂ die Gruppen -CO₂⁻, SO₃⁻ oder einen Substituenten mit der biologisch aktiven Gruppierung darstellen;

vorausgesetzt, daß ein R, R_1 , R_2 , R_3 und R_4 einen Substituenten mit der biologisch aktiven Gruppierung und mindestens ein verbleibendes R, R_1 , R_2 , R_3 und R_4 einen Substituenten mit einer hydrophilen Gruppe darstellen;

a und b unabhängig voneinander eine ganze Zahl von 0 bis 4 bedeuten, vorausgesetzt, daß die Summe von a + b eine ganze Zahl von 1 bis 4 darstellt;

m und n jeweils 0 oder 1 bedeuten, vorausgesetzt, daß ein m und n = 0 und das andere = 1 bedeutet; p und q jeweils 0 oder 1 bedeuten, vorausgesetzt, daß ein p und q = 0 und das andere = 1 bedeutet; A ein Gegenion oder die Gegenionen zum Ausgleich der Gesamtladungen auf der Konjugatgruppierung darstellt; und

X = 0 oder 1 ist.

- Diagnostisches Element nach Anspruch 7, enthaltend eine Unterlage, welche mindestens eine Reagensschicht trägt.
 - Diagnostisches Element nach Anspruch 8, enthaltend den Bindungspartner der im Konjugat vorliegenden biologisch aktiven Gruppierung.

55

35

40

45

10. Assay-Element nach Anspruch 7, worin das Konjugat durch die Formel

5

10

15

20

25

50

55

definiert ist, worin R_{12} , R_{13} , R_{14} und R_{15} unabhängig voneinander Wasserstoff, einen Substituenten mit einer hydrophilen Gruppe oder einen Substituenten mit der biologisch aktiven Gruppierung darstellen, vorausgesetzt, daß ein R_2 , R_3 , R_{12} , R_{13} , R_{14} und R_{15} einen Substituenten mit der biologisch aktiven Gruppierung und mindestens eine verbleibende Gruppe R_3 , R_{12} , R_{13} , R_{14} und R_{15} einen Substituenten mit einer hydrophilen Gruppe darstellt.

11. Assay-Element nach Anspruch 7, worin das Konjugat durch die Formel

30
$$R_{12} \longrightarrow R_{14} \longrightarrow R_{15}$$

$$R_{15} \longrightarrow R_{13}$$

$$R_{10} \longrightarrow R_{12} \longrightarrow R_{2}$$

$$R_{10} \longrightarrow R_{2}$$

$$R_{3} \longrightarrow R_{3}$$

$$R_{10} \longrightarrow R_{2}$$

$$R_{3} \longrightarrow R_{3}$$

definiert ist, worin R_{12} , R_{13} , R_{14} und R_{15} jeweils unabhängig voneinander Wasserstoff, einen Substituenten mit einer hydrophilen Gruppe oder einen Substituenten mit der biologisch aktiven Gruppierung darstellen, vorausgesetzt, daß ein R_2 , R_3 , R_{12} , R_{13} , R_{14} und R_{15} einen Substituenten mit der biologisch aktiven Gruppierung und mindestens eine verbleibende Gruppe R_3 , R_{12} , R_{13} , R_{14} und R_{15} einen Substituenten mit einer hydrophilen Gruppe darstellt.

- 12. Assay-Element nach Anspruch 10 oder 11, worin R₂ einen Substituenten mit der biologisch aktiven Gruppierung darstellt.
- Assay-Element nach einem der Ansprüche 10 oder 11, worin R₁₂ und/oder R₁₃ einen Substituenten mit einer hydrophilen Gruppe darstellen.

- 14. Assay-Element nach Anspruch 13, worin die hydrophile Gruppe eine Carbonsäure-, eine Polyether-, eine Polyalkohol-, eine Amino-, eine Sulfonsäure-, eine Phosphonsäure-, eine Phosphonsäure- oder eine Borinsäuregruppe darstellt.
- 15. Verwendung des fluoreszierenden Konjugats nach einem der Ansprüche 1 bis 3, zur Fluoreszenzfärbung von Zellen oder deren Nachweis oder Abtrennung in einer Sortiervorrichtung für fluoreszierende aktivierte Zellen.

10 Revendications

15

Revendications pour les Etats contractants suivants : AT, BE, CH, DE, FR, GB, IT, LI, LU, NL, SE

 Test diagnostic par fluorescence, comprenant le fait de combiner, dans un milieu d'essai, un échantillon suspecté de contenir un analyte, un partenaire de liaison pour cet analyte, et un conjugué de cet analyte et d'un composé fluorescent, ledit conjugué étant représenté par la formule

dans laquelle W, X, Y et Z représentent chacun, indépendamment, un groupe - $(CH_2)_a$ - (CHR_4) - $(CH_2)_b$ -, R, R₁, R₃ et R₄ représentent chacun, indépendamment, un atome d'hydrogène, un substituant contenant un groupe hydrophile ou un substituant contenant un fragment biologiquement actif,

R₂ représente -CO₂⁻, -SO₃⁻ ou un substituant contenant un fragment biologiquement actif, à condition que l'un des R, R₁, R₂, R₃ et R₄ représente un substituant contenant un fragment biologiquement actif et qu'au moins l'un des R, R₄, R₅ et R₅, restants représente un substituent

biologiquement actif et qu'au moins l'un des R, R₁, R₃ et R₄ restants représente un substituant contenant un groupe hydrophile,

a et b représentent chacun, indépendamment, un nombre entier valant de 0 à 4, à condition que la somme a + b soit un entier valant de 1 à 4,

m et n valent chacun 0 ou 1, à condition que l'un d'eux vaille 0 et l'autre 1,

p et q valent chacun 0 ou 1, à condition que l'un d'eux vaille 0 et l'autre 1,

A représente un contre-ion ou les contre-ions nécessaires pour équilibrer l'ensemble des charges présentes dans le fragment conjugué,

et X vaut 0 ou 1;

le fait de séparer le conjugué lié d'avec le conjugué libre ;

la mesure du niveau du signal de fluorescence fourni par le conjugué libre ou par le conjugué lié; et la mise en relation du niveau du signal avec la quantité dudit analyte dans l'échantillon.

50

35

40

2. Test conforme à la revendication 1, dans lequel ledit conjugué est représente par la formule

20

25

50

dans laquelle R₁₂, R₁₃, R₁₄ et R₁₅ représentent chacun, indépendamment, un atome d'hydrogène, un substituant contenant un groupe hydrophile ou un substituant contenant un fragment biologiquement actif, à condition que l'un des R₂, R₃, R₁₂, R₁₃, R₁₄ et R₁₅ représente un substituant contenant un fragment biologiquement actif, et qu'au moins l'un des R₃, R₁₂, R₁₃, R₁₄ et R₁₅ restants représente un substituant contenant un groupe hydrophile.

3. Test conforme à la revendication 1, dans lequel ledit conjugué est représenté par la formule

dans laquelle R₁₂, R₁₃, R₁₄ et R₁₅ représentent chacun, indépendamment, un atome d'hydrogène, un substituant contenant un groupe hydrophile ou un substituant contenant un fragment biologiquement actif, à condition que l'un des R₂, R₃, R₁₂, R₁₃, R₁₄ et R₁₅ représente un substituant contenant un fragment biologiquement actif, et qu'au moins l'un des R₃, R₁₂, R₁₃, R₁₄ et R₁₅ restants représente un substituant contenant un groupe hydrophile.

 Test conforme à la revendication 2 ou 3, dans lequel R₂ représente un substituant contenant un fragment biologiquement actif.

- Test conforme à l'une quelconque des revendications 2 et 3, dans lequel R₁₂ et/ou R₁₃ r présentent un substituant contenant un groupe hydrophile.
- 6. Test conforme à la revendication 5, dans lequel ledit groupe hydrophile est un groupe acide carboxylique, polyéther, polyalcool, amino, acide sulfonique, acide phosphonique, ester phosphate, acide phosphinique, acide boronique ou acide borinique.
- 7. Elément pour test diagnostic, adapté pour recevoir un échantillon d'un fluide biologique et pour fournir un signal détectable, en fonction de la présence éventuelle d'un analyte dans ledit fluide, ledit élément pour test contenant un conjugué fluorescent représenté par la formule

dans laquelle W, X, Y et Z représentent chacun, indépendamment, un groupe -(CH₂)_a-(CHR₄)-(CH₂)_b-, R, R₁, R₃ et R₄ représentent chacun, indépendamment, un atome d'hydrogène, un substituant contenant un groupe hydrophile ou un substituant contenant un fragment biologiquement actif choisi parmi un antigène, un anticorps, un haptène, un fragment Fab et une sonde d'ADN,

R₂ représente -CO₂⁻⁻ -SO₃⁻⁻ ou un substituant contenant lefit fragment biologiquement actif,

à condition que l'un des R, R₁, R₂, R₃ et R₄ représente un substituant contenant ledit fragment biologiquement actif et qu'au moins l'un des R, R₁, R₃ et R₄ restants représente un substituant contenant un groupe hydrophile,

a et b représentent chacun, indépendamment, un nombre entier valant de 0 à 4, à condition que la somme a + b soit un entier valant de 1 à 4,

m et n valent chacun 0 ou 1, à condition que l'un d'eux vaille 0 et l'autre 1,

p et q valent chacun 0 ou 1, à condition que l'un d'eux vaille 0 et l'autre 1,

A représente un contre-ion ou les contre-ions nécessaires pour équilibrer l'ensemble des charges présentes dans le fragment conjugué, et X vaut 0 ou 1.

- 8. Elément pour diagnostic, conforme à la revendication 7, qui contient un support portant au moins une couche de réactif.
 - 9. Elément pour diagnostic, conforme à la revendication 8, qui contient le partenaire de liaison dudit fragment biologiquement actif présent dans ledit conjugué.
- 50 10. Elément pour test, conforme à la revendication 7, dans lequel ledit conjugué est représenté par la formule

30

35

20

25

50

dans laquelle R₁₂, R₁₃, R₁₄ et R₁₅ représentent chacun, indépendamment, un atome d'hydrogène, un substituant contenant un groupe hydrophile ou un substituant contenant lefit fragment biologiquement actif, à condition que l'un des R₂, R₃, R₁₂, R₁₃, R₁₄ et R₁₅ représente un substituant contenant ledit fragment biologiquement actif, et qu'au moins l'un des R₃, R₁₂, R₁₃, R₁₄ et R₁₅ restants représente un substituant contenant un groupe hydrophile.

11. Elément pour test, conforme à la revendication 7, dans lequel ledit conjugué est représenté par la formule

35
$$R_{12}$$

$$R_{14}$$

$$R_{15}$$

$$R_{18}$$

$$R_{19}$$

dans laquelle R₁₂, R₁₃, R₁₄ et R₁₅ représentent chacun, indépendamment, un atome d'hydrogène, un substituant contenant un groupe hydrophile ou un substituant contenant ledit fragment biologiquement actif, à condition que l'un des R₂, R₃, R₁₂, R₁₃, R₁₄ et R₁₅ représente un substituant contenant ledit fragment biologiquement actif, et qu'au moins l'un des R₃, R₁₂, R₁₃, R₁₄ et R₁₅ restants représente un substituant contenant un groupe hydrophile.

12. Elément pour test, conforme à la revendication 10 ou 11, dans lequel R₂ représente un substituant contenant ledit fragment biologiquement actif.

EP 0 285 179 B1

- 13. Elément pour test, conforme à l'une qu lconque des r vendications 10 et 11, dans lequel R₁₂ et/ou R₁₃ représentent un substituant contenant un groupe hydrophile.
- 14. Elément pour test, conforme à la revendication 13, dans lequel ledit groupe hydrophile est un groupe acide carboxylique, polyéther, polyalcool, amino, acide sulfonique, acide phosphonique, ester phosphonique, phosphate, ester phosphate, acide phosphinique, acide boronique ou acide borinique.
- 15. Conjugué fluorescent, représenté par la formule

5

25

30

35

dans laquelle W, X, Y et Z représentent chacun, indépendamment, un groupe -(CH₂)_a-(CHR₄)-(CH₂)_b-, R, R₁, R₃ et R₄ représentent chacun, indépendamment, un atome d'hydrogène, un substituant contenant un groupe hydrophile ou un substituant contenant un fragment biologiquement actif choisi parmi un antigène, un anticorps, un haptène, un fragment Fab et une sonde d'ADN,

 R_2 représente $-CO_2^-$, $-SO_3^-$ ou un substituant contenant ledit fragment biologiquement actif, à condition que l'un des R, R₁, R₂, R₃ et R₄ représente un substituant contenant ledit fragment biologiquement actif et qu'au moins l'un des R, R₁, R₃ et R₄ restants représente un substituant contenant un groupe hydrophile,

a et b représentent chacun, indépendamment, un nombre entier valant de 0 à 4, à condition que la somme a + b soit un entier valant de 1 à 4,

m et n valent chacun 0 ou 1, à condition que l'un d'eux vaille 0 et l'autre 1,

p et q valent chacun 0 ou 1, à condition que l'un d'eux vaille 0 et l'autre 1,

A représente un contre-ion ou les contre-ions nécessaires pour équilibrer l'ensemble des charges présentes dans le fragment conjugué,

40 et X vaut 0 ou 1.

16. Conjugué fluorescent conform à la revendication 15, représenté par la formul

5

10

15

20

25

50

dans laquelle R₁₂, R₁₃, R₁₄ et R₁₅ représentent chacun, indépendamment, un atome d'hydrogène, un substituant contenant un groupe hydrophile ou un substituant contenant ledit fragment biologiquement actif, à condition que l'un des R₂, R₃, R₁₂, R₁₃, R₁₄ et R₁₅ représente un substituant contenant un fragment biologiquement actif, et qu'au moins l'un des R₃, R₁₂, R₁₃, R₁₄ et R₁₅ restants représente un substituant contenant un groupe hydrophile.

17. Conjugué fluorescent conforme à la revendication 15, représenté par la formule

dans laquelle R₁₂, R₁₃, R₁₄ et R₁₅ représentent chacun, indépendamment, un atome d'hydrogène, un substituant contenant un groupe hydrophile ou un substituant contenant ledit fragment biologiquement actif, à condition que l'un des R₂, R₃, R₁₂, R₁₃, R₁₄ et R₁₅ représente un substituant contenant ledit fragment biologiquement actif, et qu'au moins l'un des R₃, R₁₂, R₁₃, R₁₄ et R₁₅ restants représente un substituant contenant un groupe hydrophile.

5 18. Conjugué fluorescent conforme à l'une quelconque des revendications 16 et 17, dans lequel R2 représente un substituant contenant ledit fragment biologiquement actif.

19. Utilisation d'un conjugué fluorescent conform à l'une quelconque des revendications 15 à 18, pour la coloration de cellules par fluorescence, ou pour l'élimination ou la séparation dans un trieur de cellules activé par fluorescence.

5 Revendications pour les Etats contractants suivants : ES, GR

 Test diagnostic par fluorescence, comprenant le fait de combiner, dans un milieu d'essai, un échantillon suspecté de contenir un analyte, un partenaire de liaison pour cet analyte, et un conjugué de cet analyte et d'un composé fluorescent, ledit conjugué étant représenté par la formule

25

30

35

40

10

dans laquelle W, X, Y et Z représentent chacun, indépendamment, un groupe -(CH₂)_a-(CHR₄)-(CH₂)_b-, R, R₁, R₃ et R₄ représentent chacun, indépendamment, un atome d'hydrogène, un substituant contenant un groupe hydrophile ou un substituant contenant un fragment biologiquement actif,

R₂ représente -CO₂⁻ -SO₃⁻ ou un substituant contenant un fragment biologiquement actif, à condition que l'un des R, R₁, R₂, R₃ et R₄ représente un substituant contenant un fragment biologiquement actif et qu'au moins l'un des R, R₁, R₃ et R₄ restants représente un substituant contenant un groupe hydrophile,

a et b représentent chacun, indépendamment, un nombre entier valant de 0 à 4, à condition que la somme a + b soit un entier valant de 1 à 4,

m et n valent chacun 0 ou 1, à condition que l'un d'eux vaille 0 et l'autre 1, p et q valent chacun 0 ou 1, à condition que l'un d'eux vaille 0 et l'autre 1.

A représente un contre-ion ou les contre-ions nécessaires pour équilibrer l'ensemble des charges présentes dans le fragment conjugué,

et X vaut 0 ou 1;

le fait de séparer le conjugué lié d'avec le conjugué libre ;

la mesure du niveau du signal de fluorescence fourni par le conjugué libre ou par le conjugué lié ; et la mise en relation du niveau du signal avec la quantité dudit analyte dans l'échantillon.

45

50

2. Test conforme à la revendication 1, dans lequel ledit conjugué est représenté par la formule

20

25

50

dans laquelle R₁₂, R₁₃, R₁₄ et R₁₅ représentent chacun, indépendamment, un atome d'hydrogène, un substituant contenant un groupe hydrophile ou un substituant contenant un fragment biologiquement actif, à condition que l'un des R₂, R₃, R₁₂, R₁₃, R₁₄ et R₁₅ représente un substituant contenant un fragment biologiquement actif, et qu'au moins l'un des R₃, R₁₂, R₁₃, R₁₄ et R₁₅ restants représente un substituant contenant un groupe hydrophile.

3. Test conforme à la revendication 1, dans lequel ledit conjugué est représenté par la formule

dans laquelle R₁₂, R₁₃, R₁₄ et R₁₅ représentent chacun, indépendamment, un atome d'hydrogène, un substituant contenant un groupe hydrophile ou un substituant contenant un fragment biologiquement actif, à condition que l'un des R₂, R₃, R₁₂, R₁₃, R₁₄ et R₁₅ représente un substituant contenant un fragment biologiquement actif, et qu'au moins l'un des R₃, R₁₂, R₁₃, R₁₄ et R₁₅ restants représente un substituant contenant un groupe hydrophile.

 Test conforme à la revendication 2 ou 3, dans lequel R₂ représente un substituant contenant un fragment biologiquement actif.

- Test conforme à l'une quelconque des revendications 2 et 3, dans lequel R₁₂ et/ou R₁₃ représentent un substituant contenant un groupe hydrophile.
- 6. Test conforme à la revendication 5, dans lequel ledit groupe hydrophile est un groupe acide carboxylique, polyéther, polyalcool, amino, acide sulfonique, acide phosphonique, ester phosphonique, phosphate, ester phosphate, acide phosphinique, acide boronique ou acide borinique.
- 7. Elément pour test diagnostic, adapté pour recevoir un échantillon d'un fluide biologique et pour fournir un signal détectable, en fonction de la présence éventuelle d'un analyte dans ledit fluide, ledit élément pour test contenant un conjugué fluorescent représenté par la formule

dans laquelle W, X, Y et Z représentent chacun, indépendamment, un groupe -(CH₂)_a-(CHR₄)-(CH₂)_b-, R, R₁, R₃ et R₄ représentent chacun, indépendamment, un atome d'hydrogène, un substituant contenant un groupe hydrophile ou un substituant contenant un fragment biologiquement actif choisi parmi un antigène, un anticorps, un haptène, un fragment Fab et une sonde d'ADN,

R₂ représente -CO₂---SO₃- ou un substituant contenant lefit fragment biologiquement actif,

à condition que l'un des R, R₁, R₂, R₃ et R₄ représente un substituant contenant ledit fragment biologiquement actif et qu'au moins l'un des R, R₁, R₃ et R₄ restants représente un substituant contenant un groupe hydrophile,

a et b représentent chacun, indépendamment, un nombre entier valant de 0 à 4, à condition que la somme a + b soit un entier valant de 1 à 4,

m et n valent chacun 0 ou 1, à condition que l'un d'eux vaille 0 et l'autre 1,

p et q valent chacun 0 ou 1, à condition que l'un d'eux vaille 0 et l'autre 1,

A représente un contre-ion ou les contre-ions nécessaires pour équilibrer l'ensemble des charges présentes dans le fragment conjugué, et X vaut 0 ou 1.

- 8. Elément pour diagnostic, conforme à la revendication 7, qui contient un support portant au moins une couche de réactif.
 - 9. Elément pour diagnostic, conforme à la revendication 8, qui contient le partenaire de liaison dudit fragment biologiquement actif présent dans ledit conjugué.
- 50 10. Elément pour test, conforme à la revendication 7, dans lequel ledit conjugué est représenté par la formule

55

30

35

dans laquelle R₁₂, R₁₃, R₁₄ et R₁₅ représentent chacun, indépendamment, un atome d'hydrogène, un substituant contenant un groupe hydrophile ou un substituant contenant ledit fragment biologiquement actif, à condition que l'un des R₂, R₃, R₁₂, R₁₃, R₁₄ et R₁₅ représente un substituant contenant ledit fragment biologiquement actif, et qu'au moins l'un des R₃, R₁₂, R₁₃, R₁₄ et R₁₅ restants représente un substituant contenant un groupe hydrophile.

25

50

55

11. Elément pour test, conforme à la revendication 7, dans lequel ledit conjugué est représenté par la formule

dans laquelle R_{12} , R_{13} , R_{14} et R_{15} représentent chacun, indépendamment, un atome d'hydrogène, un substituant contenant un groupe hydrophile ou un substituant contenant ledit fragment biologiquement actif, à condition que l'un des R_2 , R_3 , R_{12} , R_{13} , R_{14} et R_{15} représente un substituant contenant ledit fragment biologiquement actif, et qu'au moins l'un des R_3 , R_{12} , R_{13} , R_{14} et R_{15} restants représente un substituant contenant un groupe hydrophile.

- 12. Elément pour test, conforme à la revendication 10 ou 11, dans lequel R₂ représente un substituant contenant ledit fragment biologiquement actif.
- 13. Elément pour test, conforme à l'une quelconqu des revendications 10 et 11, dans lequel R₁₂ et/ou R₁₃ représentent un substituant contenant un groupe hydrophile.

EP 0 285 179 B1

- 14. Elément pour test, conforme à la revendication 13, dans lequel ledit groupe hydrophile est un groupe acide carboxylique, polyéther, polyalcool, amino, acide sulfonique, acide phosphonique, ester phosphonique, phosphate, ester phosphate, acide phosphinique, acide boronique ou acid borinique.
- 15. Utilisation d'un conjugué fluorescent conforme à l'une quelconque des revendications 1 à 3, pour la coloration de cellules par fluorescence, ou pour l'élimination ou la séparation dans un trieur de cellules activé par fluorescence.